

# Targeting c-Myc-activated genes with a correlation method: Detection of global changes in large gene expression network dynamics

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This work studies the dynamics of a gene expression time series network. The network, which is obtained from the correlation of gene expressions, exhibits global dynamic properties that emerge after a cell state perturbation. The main features of this network appear to be more robust when compared with those obtained with a network obtained from a linear Markov model. In particular, the network properties strongly depend on the exact time sequence relationships between genes and are destroyed by random temporal data shuffling. We discuss in detail the problem of finding targets of the *c-myc* protooncogene, which encodes a transcriptional regulator whose inappropriate expression has been correlated with a wide array of malignancies. The data used for network construction are a time series of gene expression, collected by microarray analysis of a rat fibroblast cell line expressing a conditional Myc-estrogen receptor oncoprotein. We show that the correlation-based model can establish a clear relationship between network structure and the cascade of *c-myc*-activated genes.

complex systems | time series | gene interaction

The availability in modern molecular biology of methods capable of measuring the activity of thousands of genes at the same time poses the challenge of analysis and modeling of complex biological networks with thousands of units. Microarray technology is producing data on the activity of significant portions of the genome in a wide variety of cells and organisms up to the level of the entire human genome. Several techniques have been proposed to analyze the high dimensional data resulting from these experiments. Artificial neural networks, phylogenetic-type trees, clustering algorithms, and kernel methods are just a few examples (1–6).

Complex network theory has been used to characterize topological features of many biological systems such as metabolic pathways, protein–protein interactions, and neural networks (7, 8). The application of network theory to gene expression data has been not fully investigated, particularly regarding the time-dependent relationships between genes occurring while their expression level changes.

One of the key points of the network approach is the definition of the links between its elements (nodes), namely, the gene interactions from which all of the network properties are obtained. Recently, several methods for links assessment have been proposed, such as linear Markov model (LMM)-based methods (9, 10) or correlation-based methods (11–13). We choose to define links on the basis of the time correlation properties of gene expression measurements.

In this article, we show that correlation properties of gene expression time series measurements reflect very broad changes in genomic activity. The problem that we address is characterizing the gene transregulation cascade in response to *c-Myc* protooncogene activation. *C-myc* encodes a transcriptional reg-

ulator whose inappropriate expression is correlated with a wide array of malignancies. At the cellular level *c-Myc* activity has been linked with cell division, accumulation of mass, differentiation, and programmed cell death. Although the positive influence of *c-Myc* on proliferation has been appreciated for a long time, the molecular mechanisms by which these end points are achieved are not well understood. It is now clear that Myc can directly influence the expression of thousands of genes with diverse functions. A significant challenge is to integrate this wealth of information into mechanistic models that explain the biological functions of *c-Myc*. This endeavor has been greatly complicated not only by the large number of targets, but also by the weak transcriptional effects exerted by *c-Myc*. Thus, the biologically relevant downstream effectors remain to be comprehensively delineated.

The correlation method is more sensitive to the temporal structure of the data than LMM and leads to biologically relevant gene identification that is not obtained by either Markov modeling or significance analysis based only on ANOVA.

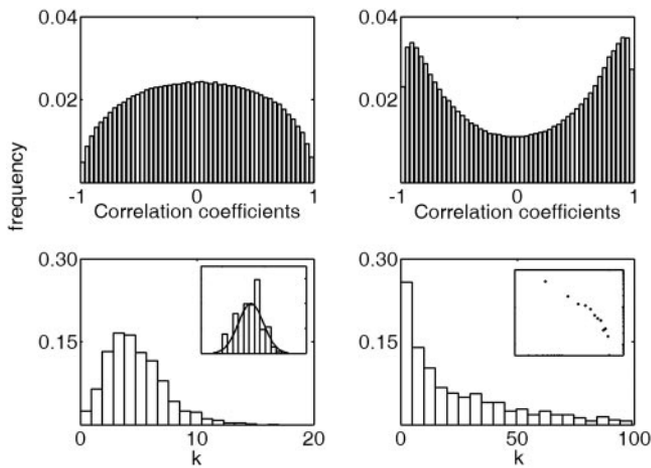
## Methods

**Gene Expression Time Series.** Two data sets of gene expression were obtained from a set of microarray experiments using genetically engineered rat cell lines. As described (ref. 14 and references therein), parental Rat-1 fibroblasts were modified by homologous recombination to knock out both copies of the *c-myc* gene (*c-myc*<sup>-/-</sup> cells). This cell line was subsequently reconstituted with a cDNA encoding a fusion protein of *c-Myc* and the human estrogen receptor (MycER). The fusion protein is synthesized continuously in the cells, but is biologically inactive in the absence of a specific ligand, 4-hydroxy tamoxifen. Binding of tamoxifen to the estrogen receptor domain elicits a conformational change that allows the fusion protein to migrate to the nucleus and act as a transcription factor. A large volume of data from several laboratories indicates that the biological activities of native *c-Myc* protein and the MycER fusion protein are similar, if not identical. Randomly cycling, exponential-phase cultures were used, and conditions were developed such that cells experienced a constant environment and were in a balanced, steady state of growth for significant periods of time. Two data sets were obtained. The first data set (*N* data set) contains the gene expression data of the *c-myc*<sup>-/-</sup> MycER cell line treated with vehicle (ethanol) only. The second data set (*T* data set) contains the gene expression data collected after the addition of tamoxifen. Samples were harvested at five time points after the addition of tamoxifen to the culture medium: 1, 2, 4, 8, and 16 h. The entire experiment was repeated on three separate occasions,

Abbreviation: LMM, linear Markov model.

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**Fig. 1.** Correlation applied to  $N$  and  $T$  data sets. (Upper) Distribution of the correlation coefficients for the subset of 1,191 probe sets selected by two-way ANOVA for the  $N$  data set (Left) and the  $T$  data set (Right). (Lower Left) Histogram of  $p(k)$  for the network obtained from the  $N$  data set. (Inset) A log normal plot of the same distribution fitted with a Gaussian distribution of same mean and variance. (Lower Right) Histogram of  $p(k)$  for the  $T$  data set. (Inset) A log-log plot of the same distribution.

providing three independent measurements for each gene and each time point. Expression profiling was done by using the Affymetrix (Santa Clara, CA) platform and U34A GeneChips (8,799 probe sets; Affymetrix).

**Significance Analysis and Data Preprocessing.** A two-way full factorial ANOVA was applied to each of the 8,799 probe sets to identify those that significantly changed expression level in time between the two conditions (data set  $N$  versus data set  $T$ ). The significance analysis was based on the general linear model that describes changes in gene expression level  $\gamma$  from the global mean  $\mu$  as caused by the combination of: changes in treatment ( $\beta$ ), i.e., database  $N$  versus database  $T$ ; changes in time ( $\alpha$ ); interaction between time and treatment ( $\gamma$ ); plus some random effects ( $\varepsilon$ ):

$$y_{ijk} = \mu + \alpha_j + \beta_i + \gamma_{ij} + \varepsilon_{ijk}, \quad [1]$$

where the index  $i$  refers to the data set ( $N$  or  $T$ ); the index  $j$  refers to time ( $j = 1, 2, 4, 8, \text{ or } 16 \text{ h}$ ); and the index  $k$  refers to the replication of the experiment for a fixed condition and time ( $k = \text{experiment } 1, 2, \text{ or } 3$ ). Probe sets with a  $P$  value corresponding to the  $\beta$  factor  $< 0.05$  were considered to be significantly affected by the treatment (i.e., activation of c-myc by tamoxifen). A total of 1,191 genes were selected with this criterion. This subset of selected probes enhances the differences between the  $N$  and  $T$  networks that we observe, but the results are similar even if considering the whole data set.

The gene expression values used for the analysis were obtained by averaging over the three experiments ( $y_{ij} = 1/3 \sum_k y_{ijk}$ ) to reduce the effects of noise in the expression level measurements.

**Network Construction: Correlation-Based Model.** In the correlation-based model, the similarity measure for the expression dynamics of two genes within the same data set is given by the correlation between the two expression-level time series. Hence, for a given data set, if  $x_{ij}$  is the expression level of a gene with label  $l$  at time  $j$ , then the similarity between two genes with labels  $l$  and  $r$ , respectively, is given by:

$$c_{lr} = \frac{\sum_j (x_{lj} - \mu_l)(x_{rj} - \mu_r)}{\sigma_l \sigma_r}, \quad [2]$$

where  $\mu_l$  and  $\mu_r$  are the averages in time of the expression levels for the two genes, and  $\sigma_l$  and  $\sigma_r$  are their standard deviations. The correlation approach can be motivated by the hypothesis that genes belonging to the same activation (or inhibition) pathway should present a similar (or opposite) expression profile in time. The adjacency matrix characterizing the network was obtained by considering only the  $c_{lr}$  coefficients whose absolute value exceeded a threshold fixed between 0.95 and 0.99. The results shown in this article were obtained for a threshold equal to 0.98. These coefficients were set equal to 1, producing a symmetric adjacency matrix  $a_{lr}$ . For each gene a connectivity degree  $k$  was defined as the total number of genes it was connected to, i.e.,  $k(l) = \sum_{r \neq l} a_{lr}$ .

**Network Construction: LMM.** In the LMM, the expression level of a gene at a given time  $t_{j+1}$  is modeled as a linear combination of the expression levels of all genes at the previous time  $t_j$ . Because measurements were not performed at equally spaced times, we interpolated the time series by using a spline interpolation to generate a total of  $n = 17$  equally spaced points in time [an alternative procedure would require an optimization technique such as simulated annealing (4)]. The model can be expressed in matrix form as follows:

$$\underbrace{(\vec{x}_1 \ \vec{x}_2 \ \dots \ \vec{x}_N)}_{\vec{x}_{t+1}} = M \underbrace{(\vec{x}_0 \ \vec{x}_1 \ \dots \ \vec{x}_{N-1})}_{\vec{x}_t}, \quad [3]$$

where  $\vec{x}_j$  is a column vector of the expression levels for all genes at time  $t_j$  (the index  $i$  relative to the data set has been dropped for convenience). Because the number of genes is larger than the number of time points, Eq. 3 does not have a unique solution. A common approach is to solve it by using the Moore–Penrose generalized matrix inverse  $X_t^+$  of  $X_t$  (a unique pseudoinverse matrix obtained by including additional constraints) via its singular value decomposition (4) such that:

$$M = X_{t+1} X_t^+.$$

Because the resulting matrix  $M$  is in general not symmetric, we applied a symmetrization procedure by averaging the corresponding off-diagonal coefficients (other symmetrization techniques lead to similar results in terms of network properties). Computation of the adjacency matrix and gene connectivity from the symmetrized  $M$  matrix was performed in the same manner as in the correlation-based model, but the threshold was set as the value corresponding to the 95th percentile.

**Validation.** Time reshuffling was used to test the time sequence dependence of the results obtained by the two techniques. By randomly shuffling the time series for each gene separately, time relationships between expression levels are broken, but the mean and standard deviation for each gene are unaltered. Properties of the gene network that truly depend on the expression level

**Table 1.** Main properties of the  $N$  and  $T$  networks obtained by the correlation method

Parameters	$N$	$T$
$k_{min}$	0	0
$k_{max}$	17	99
Mean, $k$	4.53	23.44
Standard deviation, $\sigma(k)$	2.61	23.97
Skewness $\gamma(k)$	0.89	1.16
Clustering coefficient $c(k)$	0.43	0.45

**Table 2. c-Myc target genes extracted from the selected 1,191 probe sets**

GenBank	Name	Description
D13921	Acat1	Acetyl-coenzyme A acetyltransferase 1
J02752	Acox1	Acyl-coA oxidase
AA799466	Ak2	Adenylate kinase 2
M73714	Aldh3a2	Aldehyde dehydrogenase family 3, subfamily A2
M60322	Aldr1	Aldehyde reductase 1
AI177096	Aprt	Adenine phosphoribosyl transferase transferase (APRT)
U07201	Asns	Asparagine synthetase
U00926	Atp5d	ATP synthase, F1 complex, delta subunit
At4g36870	Blh2	BEL1-like homeobox 2 protein
M81681	Blvra	Biliverdin reductase A
AA859938	Bnip31	BCL2/adenovirus E1B 19-kDa-interacting protein 3-like
AI178135	C1qbp	Complement component 1, q subcomponent binding protein
L24907	Camk1	Regulator of G-protein signaling 19
U53858	Capn1	Calpain 1
U53859	Capns1	Calpain, small subunit 1
D89069	Cbr1	Carbonyl reductase 1
AA891207	Cd36l2	CD36 antigen (collagen type I receptor, thrombospondin receptor)-like 2
D26564	Cdc37	Cell division cycle 37 homolog
L11007	Cdk4	Cyclin-dependent kinase 4
AB009999	Cds1	CDP-diacylglycerol synthase
U66470	Cgref1	Cell growth regulator with EF hand domain 1
M15882	Clta	Clathrin, light polypeptide (Lca)
D28557	Csda	Cold shock domain protein A
AI008888	Cstb	Cystatin B
AJ000485	Cyln2	Cytoplasmic linker 2
U95727	Dnaja2	Dnaj (Hsp40) homolog, subfamily A, member 2
U08976	Ech1	Enoyl coenzyme A hydratase 1
D38056	Efna1	Ephrin A1
U19516	Eif2b5	Initiation factor eIF-2Be
X03362	ErbB2	v-erb-b2 oncogene homolog 2
U36482	Erp29	Endoplasmic reticulum protein 29
J04473	Fh1	Fumarate hydratase 1
AI231547	Fkbp4	FK506 binding protein 4 (59 kDa)
M81225	Fnta	Farnesyltransferase, CAAX box, $\alpha$
AI136396	Fntb	Farnesyltransferase $\beta$ subunit
AA891857	Fxc1	Fractured callus expressed transcript 1
AA892649	Gabarap	$\gamma$ -Aminobutyric acid receptor associated protein
J03588	Gamt	Guanidinoacetate methyltransferase
D30735	Gfer	Growth factor, erv1-like
U38379	Ggh	$\gamma$ -Glutamyl hydrolase
AA944423	Gm130	cis-Golgi matrix protein GM130
AA799779	Gnpat	Acyl-CoA: dihydroxyacetone phosphate acyltransferase
U62940	Grpel1	GrpE-like 1, mitochondrial
X04229	Gstm1	Glutathione S-transferase, $\mu$ 1

**Table 2. (continued)**

GenBank	Name	Description
AB008807	Gsto1	Glutathione S-transferase $\omega$ 1
D16478	Hadha	Hydroxyacyl-Coenzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/enoyl-coenzyme A hydratase (trifunctional protein), $\alpha$ subunit
AA892036	Hdac6	Histone deacetylase 6
X52625	Hmgcs1	3-Hydroxy-3-methylglutaryl-coenzyme A synthase 1
D14048	Hnrpu	System N1 Na <sup>+</sup> and H <sup>+</sup> -coupled glutamine transporter
S57565	Hrh2	Histamine receptor H
AA957923	Mcpt2	Mast cell protease 2
U62635	Mrp123	Mitochondrial ribosomal protein L23
AF104399	Msg1	Melanocyte-specific gene 1 protein
X93495	Mtap6	Microtubule-associated protein 6
M55017	Ncl	Nucleolin
AF045564	Ndr4	N-myc downstream regulated
AA874794	Ngfrap1	Nerve growth factor receptor associated protein 1
AA998882	Nopp140	Nucleolar phosphoprotein p130
J04943	Npm1	Nucleophosmin 1
M25804	Nr1d1	Nuclear receptor subfamily 1, group D, member 1
AB015724	Nrbf1	Nuclear receptor binding factor 1
AA800679	Ns	Nucleostemin
D13309	Nsep1	Nuclease sensitive element binding protein 1
X82445	Nudc	Nuclear distribution gene C homolog
U03416	Olfm1	Olfactomedin-related ER localized protein
U26541	Pdap1	PDGFA-associated protein 1
M80601	Pdcd2	Programmed cell death 2
S82627	Pem	Placentae and embryos oncofetal gene
AI169417	Pgam1	Phosphoglycerate mutase 1
AA998446	Pitpnb	Phosphotidylinositol transfer protein, $\beta$
X71898	Plaur	Plasminogen activator, urokinase receptor
L25331	Plod	Procollagen-lysine hydroxylase
S55427	Pmp22	Peripheral myelin protein 22
AJ222691	Pold1	DNA polymerase delta, catalytic subunit
AB017711	Polr2f	Polymerase II
Z71925	Polr2g	RNA polymerase II polypeptide G
AA892298	Ppil3	Peptidylprolyl isomerase (cyclophilin)-like 3
Y17295	Prdx6	Peroxiredoxin 6
D85435	Prkcdp	PKC-delta binding protein
D26180	Prkcl1	Protein kinase C-like 1
AA891871	Prpsap1	Phosphoribosylpyrophosphate synthetase-associated protein
D10756	Psma5	Proteasome subunit, $\alpha$ type 5
D10755	Psma6	Proteasome subunit, $\alpha$ type 6
U03388	Ptgs1	Prostaglandin-endoperoxide synthase 1
L27843	Ptp4a1	Protein tyrosine phosphatase 4a1
U53475	Rab8b	GTPase Rab8b
AA956332	Rabep1	Rabaptin 5
L19699	Ralb	v-ral oncogene homolog B
U82591	Rcl	Chromosome 6 open reading frame 108

dynamics should be significantly affected by a random shuffling in time.

## Results

When c-Myc is activated by tamoxifen stimulation, the activity profile of the probe sets clearly changes into a strongly correlated regime. These findings are reflected in the histograms of the correlation coefficients for the *N* and *T* data sets (Fig. 1) and in

the main parameters of the connectivity distributions obtained from the corresponding adjacency matrices (Table 1). For the *T* data set, the number of coefficients close to +1 or -1 increases significantly. This finding is an indication that many of the 1,191 genes that were affected mostly by tamoxifen stimulation in their expression levels over time became either strongly correlated or anticorrelated.

Both networks appear to be highly clustered (Table 1), as compared with a random network with the same number of

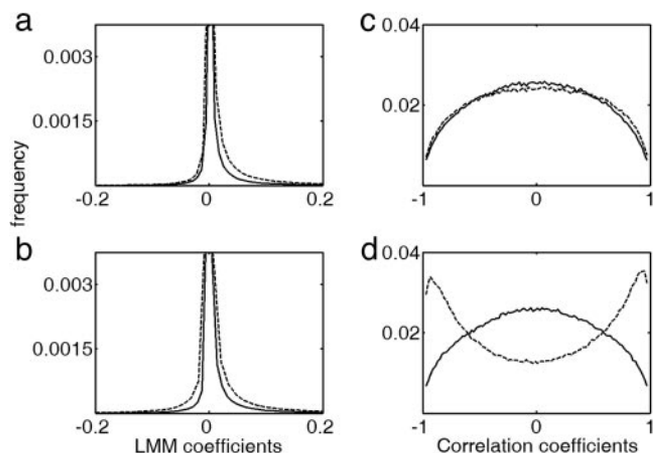
**Table 2. (continued)**

GenBank	Name	Description
X62528	Rnh1	Ribonuclease/angiogenin inhibitor
X78327	Rpl13	Ribosomal protein L13
X78167	Rpl15	Ribosomal protein L15
M20156	Rpl18	Ribosomal protein L18
M17419	Rpl5	Ribosomal protein L5
X62145	Rpl8	Ribosomal protein L8
X53377	Rps7	Ribosomal protein S7
AB002406	Ruvbl1	RuvB-like protein 1, TIP49
AA799614	Sirt2	Sirtuin 2 (SIRT2 homolog)
D12771	Slc25a5	Solute carrier family 25, adenine nucleotide translocator, member 5
AF015305	Slc29a2	Solute carrier family 29, member 2
U60882	Hrmtl12	Heterogeneous nuclear ribonucleoprotein methyltransferase-like 2
U68562	Hsp60	Heat shock protein 60 (liver)
M86389	Hspb1	Heat shock 27-kDa protein 1
U68562	Hspe1	Heat shock 10-kDa protein 1
X65036	Itga7	Integrin $\alpha$ 7
X17163	Jun	v-jun sarcoma virus oncogene homolog
M75148	Klc1	Kinesin light chain 1
M19647	Klk7	Kallikrein 7
L38644	Kpnb1	Karyopherin $\beta$ 1
D90211	Lamp2	Lysosomal membrane glycoprotein 2
U19614	Lap1c	Lamina-associated polypeptide 1C
M69055	Lgfbp6	Insulin-like growth factor binding protein 6
A1234060	Lox	Lysyl oxidase
M61177	Mapk3	Mitogen-activated protein kinase, ERK1
AA899253	Marcks	Myristoylated alanine-rich protein kinase C substrate
AI011498	Smarcd2	SWI/SNF-related, matrix-associated, actin-dependent regulator of chromatin, subfamily d, member 2
AF007758	Snca	Synuclein, $\alpha$
AI030175	Sord	Sorbitol dehydrogenase
D37920	Sqle	Squalene epoxidase
J05035	Srd5a1	Steroid 5 $\alpha$ -reductase 1
Y15068	Stip1	Stress-induced-phosphoprotein 1 (Hsp70/Hsp90-organizing protein)
D12927	Tcea2	Transcription elongation factor A2
M58040	Tfrc	Transferrin receptor
M61142	Thop1	Thimet oligopeptidase 1
AB006451	Timm23	Translocase of inner mitochondrial membrane 23 homolog
U09256	Tkt	Transketolase
S63830	Vamp3	Vesicle-associated membrane protein 3
U14746	Vhl	von Hippel-Lindau syndrome homolog
AA875455	Wig1	p53-Activated gene 608
U96490	Yif1p	Yip1p-interacting factor
S55223	Ywhab	Tyrosine 3-monooxygenase, tryptophan 5-monooxygenase activation protein, $\beta$ polypeptide

Probes were chosen as those that mostly changed their connectivity degree between the *N* and *T* data sets.

nodes and average connectivity degree. The *T* connectivity degree distribution is much more broad and skewed, whereas the *N* connectivity degree distribution is peaked around its average value.

Considering the change in connectivity as an index for ranking the involvement of a gene in the c-Myc activation cascade, we looked at the distribution of the differences in connectivity of the probe sets (Table 2 shows a list of genes extracted from the upper tail of such a distribution). Application of a random permutation



**Fig. 2.** Effects of data reshuffling on LMM and correlation coefficients distribution obtained from the 1,191 selected probe sets. Dashed lines indicate original data. Solid lines indicate reshuffled data. (a) *N* data set, LMM. (b) *T* data set, LMM. (c) *N* data set, correlation. (d) *T* data set, correlation.

to the time series confirmed that this results depend on the exact time ordering of the gene expression levels (Fig. 2). Some features of the network structure, like the assortative mixing property (15) and the differences between the *N* and *T* data sets, are completely disrupted by time shuffling, leading to networks very similar to those obtained starting from randomly generated data of the same dimensionality, mean, and variance (data not shown).

In comparison, the gene network constructed with the LMM appears to be completely insensitive to the effects of tamoxifen. The  $p(k)$  distribution for *N* and *T* follows a power-law function  $p(k) \propto k^{-\alpha}$  with a very similar exponent,  $\alpha_N = 2.41 \pm 0.16$  and  $\alpha_T = 2.41 \pm 0.12$ , respectively. There is no evident change between the *T* and *N* networks. Moreover, the main properties of the *N* and *T* networks, namely the power-law exponent and disassortative mixing property (15), are left unchanged by time reshuffling. Thus, even if the individual genes connectivity degree changes from the *N* and *T* data sets, the insensitivity to time shuffling casts some doubts on the reliability and significance of such changes in the LMM.

### Discussion

A correlation-based model was used to identify a gene interaction network, based on a time series of gene expression measurements, resulting from the acute activation of an engineered c-Myc transcription factor in a c-myc null cell line. The global properties of the resulting network were strongly affected by c-Myc activation. The comparison between the networks obtained with the different data sets led to the identification of unique c-Myc targets. The list of genes found with this method contains some of the genes found in ref. 14 but also contains many genes that were not found before to our knowledge, pointing to the possibility that the potential list of c-Myc targets may be much larger than what was previously observed.

These network properties were disrupted by time reshuffling of the data, confirming the hypothesis that they refer to real information contained in gene expression dynamics.

The same analysis was performed on the gene network obtained with a LMM, which has been proposed in the past for the analysis of time-dependent genomic measurements. The global features of this network did not significantly change neither in response to c-Myc activation, nor after time reshuffling of the data, suggesting that they depend on some global properties of the data set distribution and not on the exact details of gene expression dynamics.

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